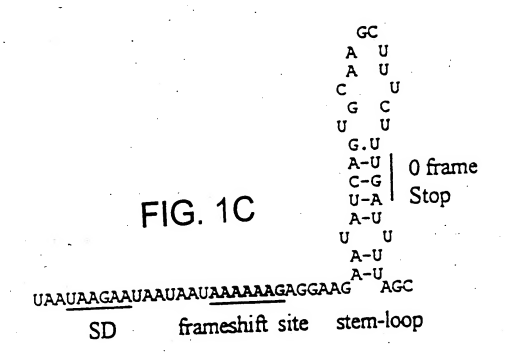


start of HP0093 ccttgggtattgactatagacgcctagacat CGCTAAAAACAGCGTGTTTGTGCATATAAGAAGAGGGGATTATGTGGGGATTGGCTGTCA CGCTAAAAACAGCGTATTTGTGCATATAAGAAGAGGGGATTATGTGGGGATTGGCTGTCA GCTTGGTATTGATTATCAAAAAAGGCGCTTGAGTATATGGCAAAGCGCGTGCCAAACAT line 3 line 4

FIG. 1B

\*\*\*stop of - 1 frame



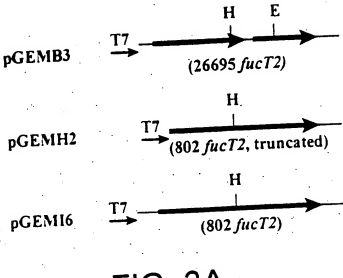


FIG. 3A

															J		
	gaac	acto	ac a	acgc	tctt	t tt	caaa	taaa	aaa	attca	aat	gatt	tgaa	ag c	gtta	cccca	60
	cttt	ttac	ggc t	ttta	ttga	a aa	aggg	jcttt	aaa	agtto	ggct	aaaa	rage	gcg t	ttta	tttga	120
	aaaa	caaa	rdd d	ggtt	ja at	g go	t tt	t aa	ia gt	g gt	g ca	ia at	t to	ge go	gg gg	g ctt	1/2
					Me	t Al	La Pr	ie r?	/5 V a	Z Va	IT CI	.n 11	re c		Ly Gi	y Leu	
	aaa	aar	caa	ato	rrr	_	tac	act	tÉc	act	aaa	agt	ttq	_	aaa	cac	220
	61 v	Asn	Gin	Met	Phe	Gln	Tvr	Ala	Phe	Ala	Lys	Ser	Leu	Gln	Lys	His	
	<b>-</b> - ,		15				- 1 -	20			•		25		-		
	ctt	aat	acg	ccc	gtg	cta	tta	gac	act	act	tct	ttt	gat	tgg	agc	aat	268
	Leu	Asn	Thr	Pro	Val	Leu	Leu	Asp	Thr	Thr	Ser	Phe	Asp	Trp	Ser	Asn	
		30					35					40					316
	agg	aaa	atg	caa	tta	gag	ctt	ttc	cct	att	gat	ttg	CCC	Tur	gcg	aat	316
	Arg 45	Lys	Met	GIn	Leu	50	reu	rne	PIO	ile	ASP 55	Leu	FIO	1 7 1	Ala	60	
		222	aaa	arc	act		act	aaa	ato	caa		ctc	ccc	aag	tta		364
	Ala	Lvs	Glu	Ile	Ala	Ile	Ala	Lvs	Met	Gln	His	Leu	Pro	Lys	Leu	Val	
					65					70					75	•	
	aga	gat	gca	ctc	aaa	tac	ata	-gga	ttt	gat	agg	gtg	agt	caa	gaa	atc	412
	Arg	Asp	Ala	Leu	Lys	Tyr	Ile	Gly		Asp	Arg	Val	Ser	Gln	Glu	Ile	
				80					85					90			160
	gtt	ttt	gaa	tac	gag	CCT	aaa	ttg	tta	aag	Dro	Sor	Ara	LEG	act Thr	Tur	460
	val	Phe	95	Tyr	GIU	Pro	Lys	100	rea.	Lys	FLO	261	105	Dea	1111	.y.	
	+++	ttt		tat	ttc	caa	gat		cga	tat	ttt	gat	-,-	ata	tcc	tct	508
	Phe	Phe	Glv	Tyr	Phe	Gln	Asp	Pro	Arg	Tyr	Phe	Asp	Ala	Ile	Ser	Ser	
	•	110					115					120					
	tta	atc	aag	caa	acc	ttc	act	cta	ccc	ccc	CCC	ccc	gaa	aat	aat	aaa	556
		Ile	Lys	Gln	Thr		Thr	Leu	Pro	Pro		Pro	Glu	Asn	Asn	Lys	
	125					130					1:35			+ = +	-+~	140	604
	aat	aat	aat	aaa	aaa	gag	gaa	Glu	Tur	Glo	Ara	Lus	Leu	Ser	ttg Leu	Tle	004
	ASN	ASI	ASI	гуs	145	GIU	Gru	Giu	ı yı	150	nr 9		204	-	155		
	tta	acc	act	aaa	aac	agc	gta	ttt	gtg	cat	ata	aga	aga	ggg	gat	tat	652
	Leu	Ála	Ála	Lys	Asn	Ser	Val	Phe	Val	His	Ile	Arg	Arg	Gly	Asp	Tyr	
				160					165					170			700
	gtg	ggg	att	ggc	tgt	cag	ctt	ggt	att.	gat	tat	Cla	aaa	aag	gcg	CCC	700
	Val	GTA	,	GLY	Cys	Gin	Leu	180	TTE	ASP	ryr	GIII	185	rys	Ala	Leu	
	~ 3 ~	<b>+</b> = + .	175	aca.	аал	cac	ara		aac	atg	gag	ctt		ata	tit	tac	748
	Glu	Tur	Met	Ala	Lvs	Ara	·Val	Pro	Asn	Met	Glu	Leu	Phe	Val	Phe	Cys	
		190					195					200					
	gaa	gac	tta	aaa	ttc.	acg	caa	aat	ctt	gat	ctt	ggc	tac	cct	ttc.	acg	796
		Asp	Leu	Lys	Phe		Gln	Asn	Leu	Asp		Gly	Tyr	Pro	Phe	Thr	
	205				1	210				~~~	215	<b>+</b> = <b>+</b>		ast.	250	220 ct.g	844
	gac	Mar	Thr	Thr	agg	Asn	Lvs	Glu	Glu	Glu	Ala	Tyr	Tro	Asp	atg Met	Leu	011
	vab		1111		225	٠.٥٢	2,3			230					235	•	
	ctc	atg	caa	t.ct	tgc	aag.	cat	ggc	att	atc	gct	aat	agc	act	tat	agc ·	892.
	Leu	Met	Gln	Ser	Cys	Lys	His	Gly	Ile	Ile	Ala	Asn	Ser	Thr	Tyr	Ser·	
			٠.	240		•			245	- 8							040
	tgg	tgg	gcg	gct	tat	ttg	atg	gaa	aac	cca	gaa	-aaa	atc	Tle	att Ile	G) v	940
	Trp	Trp	255	Ala	Tyr	Leu	Me C	260	A3II	710	GIG	Lys	265	116	. 116	O1 y	
	ccc	aaa		taa	ctt	ttt	ggg		gaa	aat	att	ctt		aag	gaa	tgg	988 -
٠	Pro	Lys	His	Trp	Leu	Phe	Gly	His	Glu	Asn.	Ile	Leu	Cys	Lys	Glu	Trp	
		270					275					280.		•			1026
	gtg	aaa	ata	gaa	tcc	cat	ttt	gag	gta	aaa	tcc.	caa	aaa	tat	aac	gct	1036
		Lys	Ile	Glu	Ser	His	Phe	Glu	Val	Lys	ser	GIN	гÀг	Tyr	Asn	300 ·	
	285					290	~rr=	craa.	a ddi	ctta.	295 atct	tta.	attt	tao .	atco	jattte	1096
		agcg: atag:						- Lay	- 99			9					1119
		acay	-ya '	4-4-			. <del>.</del>						FIC	7	10		
												ı		J.	. U		

;	Mammalian mammalian	N bacterial	Motif III FIG. 2A	II MOCIF III	VGVHVRRGDYL (215-225) GTFGFWAAYL (308-317)	VGVHVRRGDYV (196-206) GTFGIWAAYL (289-298)	VFVHIRRGDYV (163-173) STYSWWAAYL (249-258)	VGIHIRRGDFV (158-168) STFSWWAAIL (241-250)	ICVSIRRGDYV (173-183) SSFSWWTEFL (263-272)
l Catalytic	Z		Motif II Ma	Motif I Motif II	XA (87-97)	GRLGNOMGEYA (70-80) VGVHVRR	GGLGNQMFQYA (10-20) VFVHIRR	GGLGNQLFQVA (9-19) VGIHIRR	GNLGNQLFIYA (8-18) ICVSIRR
Cyt   TM <sub>1</sub> Stem			Motif I	M	man Futl (365 aa) GRFG1		Hp FucT2 (300 aa) GGLG	_	L1 EpsH (309 aa) GNLG

**UA802 wt** 

 $\Delta \mathbf{H}$ 



FIG. 4

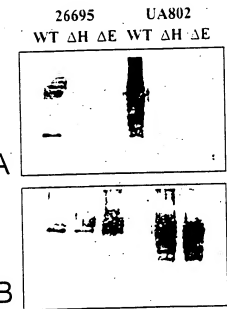


FIG. 6A

FIG. 6B

## Type 2 chain precursor (LacNAc)

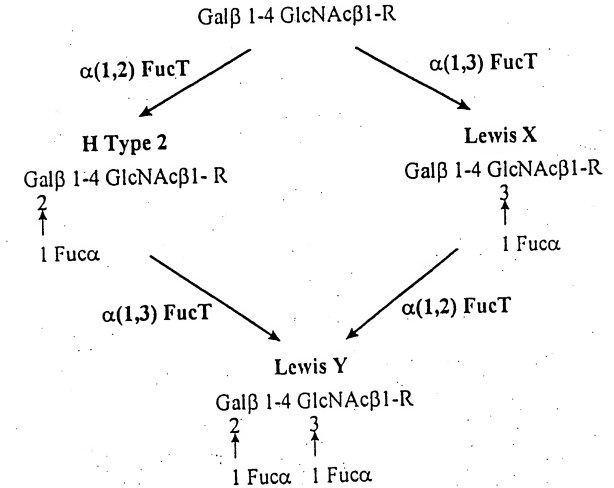


FIG. 5

